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(54) Title: PRODUCTION OF GAMMA LINOLENIC ACID BY A Δ6-DESATURASE

(57) Abstract

Linoleic acid is converted into γ -linolenic acid by the enzyme $\Delta 6$ -desaturase. The present invention is directed to an isolated nucleic acid comprising the $\Delta 6$ -desaturase gene. More particularly, the isolated nucleic acid comprises the promoter, coding region and termination regions of the $\Delta 6$ -desaturase gene. The present invention provides recombinant constructions comprising the $\Delta 6$ -desaturase coding region in functional combination with heterologous regulatory sequences. The nucleic acids and recombinant constructions of the instant invention are useful in the production of GLA in transgenic organisms.

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PRODUCTION OF GAMMA LINOLENIC ACID

BY A A6-DESATURASE

Linoleic acid (18:2) (LA) is transformed into gamma linolenic acid (18:3) (GLA) by the enzyme $\Delta 6$ -5 desaturase. When this enzyme, or the nucleic acid encoding it, is transferred into LA-producing cells, GLA is produced. The present invention provides a nucleic acid comprising the A6-desaturase gene. More specifically, the nucleic acid comprises the promoter, 10 coding region and termination regions of the $\Delta6$ desaturase gene. The present invention is further directed to recombinant constructions comprising a $\Delta 6$ desaturase coding region in functional combination with heterologous regulatory sequences. The nucleic acids 15 and recombinant constructions of the instant invention are useful in the production of GLA in transgenic organisms.

Unsaturated fatty acids such as linoleic $(C_{18}\Delta^{9,12})$ and α -linolenic $(C_{18}\Delta^{9,12,15})$ acids are 20 essential dietary constituents that cannot be synthesized by vertebrates since vertebrate cells can introduce double bonds at the Δ^9 position of fatty acids but cannot introduce additional double bonds between the Δ^9 double bond and the methyl-terminus of the fatty acid 25 chain. Because they are precursors of other products, linoleic and α -linolenic acids are essential fatty acids, and are usually obtained from plant sources. Linoleic acid can be converted by mammals into γ linolenic acid (GLA, $C_{18}\Delta^{6.9.12}$) which can in turn be converted to arachidonic acid (20:4), a critically important fatty acid since it is an essential precursor of most prostaglandins.

The dietary provision of linoleic acid, by virtue l of its resulting conversion to GLA and arachidonic acid, satisfies the dietary need for GLA and arachidonic acid. However, a relationship has been demonstrated between consumption of saturated fats and health risks such as 5 hypercholesterolemia, atherosclerosis and other chemical disorders which correlate with susceptibility to coronary disease, while the consumption of unsaturated fats has been associated with decreased blood cholesterol concentration and reduced risk of 10 atherosclerosis. The therapeutic benefits of dietary GLA may result from GLA being a precursor to arachidonic acid and thus subsequently contributing to prostaglandin synthesis. Accordingly, consumption of the more unsaturated GLA, rather than linoleic acid, has 15 potential health benefits. However, GLA is not present in virtually any commercially grown crop plant.

Linoleic acid is converted into GLA by the enzyme $\Delta 6$ -desaturase. $\Delta 6$ -desaturase, an enzyme of about 359 amino acids, has a membrane-bound domain and an active site for desaturation of fatty acids. When this enzyme is transferred into cells which endogenously produce linoleic acid but not GLA, GLA is produced. The present invention, by providing the gene encoding $\Delta 6$ -desaturase, allows the production of transgenic organisms which contain functional $\Delta 6$ -desaturase and which produce GLA. In addition to allowing production of large amounts of GLA, the present invention provides new dietary sources of GLA.

30 The present invention is directed to an isolated A6-desaturase gene. Specifically, the isolated gene

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l comprises the $\Delta 6$ -desaturase promoter, coding region, and termination region.

The present invention is further directed to expression vectors comprising the $\Delta 6$ -desaturase promoter, coding region and termination region.

The present invention is also directed to expression vectors comprising a $\Delta 6$ -desaturase coding region in functional combination with heterologous regulatory regions, i.e. elements not derived from the $\Delta 6$ -desaturase gene.

Cells and organisms comprising the vectors of the present invention, and progeny of such organisms, are also provided by the present invention.

The present invention further provides isolated

15 bacterial A6-desaturase and is still further directed to
an isolated nucleic acid encoding bacterial A6desaturase.

The present invention further provides a method for producing plants with increased gamma linolenic acid (GLA) content which comprises transforming a plant cell with an isolated nucleic acid of the present invention and regenerating a plant with increased GLA content from said plant cell.

A method for producing chilling tolerant plants is also provided by the present invention.

Fig. 1 depicts the hydropathy profiles of the deduced amino acid sequences of Synechocystis A6-desaturase (Panel A) and A12-desaturase (Panel B). Putative membrane spanning regions are indicated by solid bars. Hydrophobic index was calculated for a window size of 19 amino acid residues [Kyte, et al. (1982) J. Molec. Biol. 157].

Fig. 2 provides gas liquid chromatography profiles of wild type (Panel A) and transgenic (Panel B) Anabaena.

Fig. 3 is a diagram of maps of cosmid cSy75,

5 cSy13 and cSy7 with overlapping regions and subclones.

The origins of subclones of cSy75, cSy75-3.5 and cSy7

are indicated by the dashed diagonal lines. Restriction sites that have been inactivated are in parentheses.

Fig. 4 provides gas liquid chromatography

10 profiles of wild type (Panel A) and transgenic (Panel B)

tobacco.

The present invention provides an isolated nucleic acid encoding A6-desaturase. To identify a nucleic acid encoding $\Delta 6$ -desaturase, DNA is isolated from an organism which produces GLA. Said organism can be, for example, an animal cell, certain fungi (e.g. Mortierella), certain bacteria (e.g. Synechocystis) or certain plants (borage, Oenothera, currants). isolation of genomic DNA can be accomplished by a 20 variety of methods well-known to one of ordinary skill in the art, as exemplified by Sambrook et al. (1989) in Molecular Cloning: A Laboratory Manual, Cold Spring Harbor, NY. The isolated DNA is fragmented by physical methods or enzymatic digestion and cloned into an 25 appropriate vector, e.g. a bacteriophage or cosmid vector, by any of a variety of well-known methods which can be found in references such as Sambrook et al. (1989). Expression vectors containing the DNA of the present invention are specifically contemplated herein. 30 DNA encoding \$6-desaturase can be identified by gain of function analysis. The vector containing fragmented DNA is transferred, for example by infection,

- transconjugation, transfection, into a host organism
 that produces linoleic acid but not GLA. As used
 herein, "transformation" refers generally to the
 incorporation of foreign DNA into a host cell. Methods
- for introducing recombinant DNA into a host organism are known to one of ordinary skill in the art and can be found, for example, in Sambrook et al. (1989).

 Production of GLA by these organisms (i.e., gain of function) is assayed, for example by gas chromatography
- or other methods known to the ordinarily skilled artisan. Organisms which are induced to produce GLA, i.e. have gained function by the introduction of the vector, are identified as expressing DNA encoding A6-desaturase, and said DNA is recovered from the
- organisms. The recovered DNA can again be fragmented, cloned with expression vectors, and functionally assessed by the above procedures to define with more particularity the DNA encoding \$\(\delta \)-desaturase.

As an example of the present invention, random

DNA is isolated from the cyanobacteria Synechocystis

Pasteur Culture Collection (PCC) 6803, American Type

Culture Collection (ATCC) 27184, cloned into a cosmid

vector, and introduced by transconjugation into the GLA
deficient cyanobacterium Anabaena strain PCC 7120, ATCC

27893. Production of GLA from Anabaena linoleic acid is

monitored by gas chromatography and the corresponding

DNA fragment is isolated.

The isolated DNA is sequenced by methods well-known to one of ordinary skill in the art as found, for example, in Sambrook et al. (1989).

In accordance with the present invention, a DNA comprising a $\Delta 6$ -desaturase gene has been isolated. More

particularly, a 3.588 kilobase (kb) DNA comprising a $\Delta6$ ı desaturase gene has been isolated from the cyanobacteria Synechocystis. The nucleotide sequence of the 3.588 kb DNA was determined and is shown in SEQ ID NO:1. Open reading frames defining potential coding regions are present from nucleotide 317 to 1507 and from nucleotide 2002 to 3081. To define the nucleotides responsible for encoding A6-desaturase, the 3.588 kb fragment that confers A6-desaturase activity is cleaved into two subfragments, each of which contains only one open 10 reading frame. Fragment ORF1 contains nucleotides 1 through 1704, while fragment ORF2 contains nucleotides 1705 through 3588. Each fragment is subcloned in both forward and reverse orientations into a conjugal 15 expression vector (AM542, Wolk et al. [1984] Proc. Natl. Acad. Sci. USA 81, 1561) that contains a cyanobacterial carboxylase promoter. The resulting constructs (i.e. ORF1(F), ORF1(R), ORF2(F) and ORF2(R)] are conjugated to wild-type Anabaena PCC 7120 by standard methods (see, 20 for example, Wolk et al. (1984) Proc. Natl. Acad. Sci. USA 81, 1561). Conjugated cells of Anabaena are identified as NeoR green colonies on a brown background of dying non-conjugated cells after two weeks of growth on selective media (standard mineral media BG11N + 25 containing 30µg/ml of neomycin according to Rippka et al., (1979) J. Gen Microbiol. 111, 1). The green colonies are selected and grown in selective liquid media (BG11N + with 15µg/ml neomycin). Lipids are extracted by standard methods (e.g. Dahmer et al., 30 (1989) Journal of American Oil Chemical Society 66, 543) from the resulting transconjugants containing the

forward and reverse oriented ORF1 and ORF2 constructs.

- For comparison, lipids are also extracted from wild-type cultures of <u>Anabaena</u> and <u>Synechocystis</u>. The fatty acid methyl esters are analyzed by gas liquid chromatography (GLC), for example with a Tracor-560 gas liquid
- 5 chromatograph equipped with a hydrogen flame ionization detector and a capillary column. The results of GLC analysis are shown in Table 1.

Table 1: Occurrence of C18 fatty acids in wild-type and transgenic cyanobacteria

0	SOURCE	18:0	18:1	18:2	γ18:3	a18:3	18:4
	Anabaena (wild type)	+	+	+		+	· -
	Anabaena + ORF1(F)	+	+	+	_	+	-
5	Anabaena + ORF1(R)	+	.+	,+	-	+	-
	Anabaena + ORF2(F)	+	+	+	+	+	+
	Anabaena + ORF2(R)	+	+	+	_	+	_
	Synechocystis (wild type)	+	+	+	+	- 1	-

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As assessed by GLC analysis, GLA deficient Anabaena gain the function of GLA production when the construct containing ORF2 in forward orientation is introduced by transconjugation. Transconjugants containing constructs with ORF2 in reverse orientation to the carboxylase promoter, or ORF1 in either orientation, show no GLA production. This analysis demonstrates that the single open reading frame (ORF2) within the 1884 bp fragment encodes $\triangle 6$ -desaturase. The 1884 bp fragment is shown as SEQ ID NO:3. This is substantiated by the overall similarity of the hydropathy profiles between $\triangle 6$ -desaturase and $\triangle 12$ -

desaturase [Wada et al. (1990) Nature 347] as shown in Fig. 1 as (A) and (B), respectively.

Isolated nucleic acids encoding A6-desaturase can be identified from other GLA-producing organisms by the 5 gain of function analysis described above, or by nucleic acid hybridization techniques using the isolated nucleic acid which encodes Anabaena A6-desaturase as a hybridization probe. Both genomic and cDNA cloning methods are known to the skilled artisan and are contemplated by the present invention. 10 hybridization probe can comprise the entire DNA sequence disclosed as SEQ. ID NO:1, or a restriction fragment or other DNA fragment thereof, including an oligonucleotide probe. Methods for cloning homologous genes by cross-15 hybridization are known to the ordinarily skilled artisan and can be found, for example, in Sambrook (1989) and Beltz et al. (1983) Methods in Enzymology 100, 266.

Transgenic organisms which gain the function of

GLA production by introduction of DNA encoding Δdesaturase also gain the function of octadecatetraeonic
acid (18:4Δ^{6.9.12.15}) production. Octadecatetraeonic
acid is present normally in fish oils and in some plant
species of the Boraginaceae family (Craig et al. [1964]

J. Amer. Oil Chem. Soc. 41, 209-211; Gross et al. [1976]
Can. J. Plant Sci. 56, 659-664). In the transgenic
organisms of the present invention, octadecatetraenoic
acid results from further desaturation of α-linolenic
acid by Δ6-desaturase or desaturation of GLA by Δ15desaturase.

The 359 amino acids encoded by ORF2, i.e. the open reading frame encoding $\Delta 6$ -desaturase, are shown as

- 1 SEQ. ID NO:2. The present invention further contemplates other nucleotide sequences which encode the amino acids of SEQ ID NO:2. It is within the ken of the ordinarily skilled artisan to identify such sequences
- which result, for example, from the degeneracy of the genetic code. Furthermore, one of ordinary skill in the art can determine, by the gain of function analysis described hereinabove, smaller subfragments of the 1884 bp fragment containing ORF2 which encode A6-desaturase.

The present invention contemplates any such polypeptide fragment of Δ6-desaturase and the nucleic acids therefor which retain activity for converting LA to GLA.

In another aspect of the present invention, a

vector containing the 1884 bp fragment or a smaller
fragment containing the promoter, coding sequence and
termination region of the A6-desaturase gene is
transferred into an organism, for example,
cyanobacteria, in which the A6-desaturase promoter and
termination regions are functional. Accordingly,
organisms producing recombinant A6-desaturase are
provided by this invention. Yet another aspect of this
invention provides isolated A6-desaturase, which can be
purified from the recombinant organisms by standard
methods of protein purification. (For example, see
Ausubel et al. [1987] Current Protocols in Molecular
Biology, Green Publishing Associates, New York).

Vectors containing DNA encoding \$\triangle 6\$-desaturase are also provided by the present invention. It will be 30 apparent to one of ordinary skill in the art that appropriate vectors can be constructed to direct the expression of the \$\triangle 6\$-desaturase coding sequence in a

variety of organisms. Replicable expression vectors are 1 particularly preferred. Replicable expression vectors as described herein are DNA or RNA molecules engineered for controlled expression of a desired gene, i.e. the A6-desaturase gene. Preferably the vectors are plasmids, bacteriophages, cosmids or viruses. Shuttle vectors, e.g. as described by Wolk et al. (1984) Proc. Natl. Acad. Sci. USA, 1561-1565 and Bustos et al. (1991) J. Bacteriol. 174, 7525-7533, are also contemplated in accordance with the present invention. Sambrook et al. 10 (1989), Goeddel, ed. (1990) Methods in Enzymology 185 Academic Press, and Perbal (1988) A Practical Guide to Molecular Cloning, John Wiley and Sons, Inc., provide detailed reviews of vectors into which a nucleic acid encoding the present \$6\$-desaturase can be inserted and 15 expressed. Such vectors also contain nucleic acid sequences which can effect expression of nucleic acids encoding A6-desaturase. Sequence elements capable of effecting expression of a gene product include 20 promoters, enhancer elements, upstream activating sequences, transcription termination signals and polyadenylation sites. Both constitutive and tissue specific promoters are contemplated. For transformation of plant cells, the cauliflower mosaic virus (CaMV) 35S 25 promoter and promoters which are regulated during plant seed maturation are of particular interest. All such promoter and transcriptional regulatory elements, singly or in combination, are contemplated for use in the present replicable expression vectors and are known to 30 one of ordinary skill in the art. The CaMV 355 promoter is described, for example, by Restrepo et al. (1990)

Plant Cell 2, 987. Genetically engineered and mutated regulatory sequences are also contemplated.

The ordinarily skilled artisan can determine vectors and regulatory elements suitable for expression 5 in a particular host cell. For example, a vector comprising the promoter from the gene encoding the carboxylase of Anabaena operably linked to the coding region of A6-desaturase and further operably linked to a termination signal from Synechocystis is appropriate for 10 expression of \(\Delta 6 - \text{desaturase} \) in cyanobacteria. linked" in this context means that the promoter and terminator sequences effectively function to regulate transcription. As a further example, a vector appropriate for expression of A6-desaturase in 15 transgenic plants can comprise a seed-specific promoter sequence derived from helianthinin, napin, or glycin operably linked to the A6-desaturase coding region and further operably linked to a seed termination signal or the nopaline synthase termination signal.

In particular, the helianthinin regulatory elements disclosed in applicant's copending U.S. Application Serial No. 682,354, filed April 8, 1991 and incorporated herein by reference, are contemplated as promoter elements to direct the expression of the Δ6-25 desaturase of the present invention.

Modifications of the nucleotide sequences or regulatory elements disclosed herein which maintain the functions contemplated herein are within the scope of this invention. Such modifications include insertions, substitutions and deletions, and specifically substitutions which reflect the degeneracy of the genetic code.

1 Standard techniques for the construction of such hybrid vectors are well-known to those of ordinary skill in the art and can be found in references such as Sambrook et al. (1989), or any of the myriad of 5 laboratory manuals on recombinant DNA technology that are widely available. A variety of strategies are available for ligating fragments of DNA, the choice of which depends on the nature of the termini of the DNA fragments. It is further contemplated in accordance with the present invention to include in the hybrid 10 vectors other nucleotide sequence elements which facilitate cloning, expression or processing, for example sequences encoding signal peptides, a sequence encoding KDEL, which is required for retention of 15 proteins in the endoplasmic reticulum or sequences encoding transit peptides which direct A6-desaturase to the chloroplast. Such sequences are known to one of ordinary skill in the art. An optimized transit peptide is described, for example, by Van den Broeck et al. 20 (1985) Nature 313, 358. Prokaryotic and eukaryotic signal sequences are disclosed, for example, by Michaelis et al. (1982) Ann. Rev. Microbiol. 36, 425.

A further aspect of the instant invention provides organisms other than cyanobacteria which

25 contain the DNA encoding the \$\times 6\$-desaturase of the present invention. The transgenic organisms contemplated in accordance with the present invention include bacteria, cyanobacteria, fungi, and plants and animals. The isolated DNA of the present invention can be introduced into the host by methods known in the art, for example infection, transfection, transformation or transconjugation. Techniques for transferring the DNA

of the present invention into such organisms are widely known and provided in references such as Sambrook et al. (1989).

A variety of plant transformation methods are known. The A6-desaturase gene can be introduced into plants by a leaf disk transformation-regeneration procedure as described by Horsch et al. (1985) Science 227, 1229. Other methods of transformation, such as protoplast culture (Horsch et al. (1984) Science 223, 496; DeBlock et al. (1984) EMBO J. 2 2143. Barton et

10 496; DeBlock et al. (1984) EMBO J. 2, 2143; Barton et al. (1983) Cell 32, 1033) can also be used and are within the scope of this invention. In a preferred embodiment plants are transformed with Agrobacterium-derived vectors. However, other methods are available

to insert the \(\alpha 6 - \text{desaturase} \) gene of the present invention into plant cells. Such alternative methods include biolistic approaches (Klein et al. (1987) Nature \(\frac{327}{7}, 70 \)), electroporation, chemically-induced DNA uptake, and use of viruses or pollen as vectors.

When necessary for the transformation method, the Δ6-desaturase gene of the present invention can be inserted into a plant transformation vector, e.g. the binary vector described by Bevan (1984) Nucleic Acids Res. 12, 8111. Plant transformation vectors can be

derived by modifying the natural gene transfer system of Agrobacterium tumefaciens. The natural system comprises large Ti (tumor-inducing)-plasmids containing a large segment, known as T-DNA, which is transferred to transformed plants. Another segment of the Ti plasmid,

30 the <u>vir</u> region, is responsible for T-DNA transfer. The T-DNA region is bordered by terminal repeats. In the modified binary vectors the tumor-inducing genes have

- been deleted and the functions of the <u>vir</u> region are utilized to transfer foreign DNA bordered by the T-DNA border sequences. The T-region also contains a selectable marker for antibiotic resistance, and a
- multiple cloning site for inserting sequences for transfer. Such engineered strains are known as "disarmed" A. tumefaciens strains, and allow the efficient transformation of sequences bordered by the Tregion into the nuclear genomes of plants.
- Surface-sterilized leaf disks are inoculated with the "disarmed" foreign DNA-containing A. tumefaciens, cultured for two days, and then transferred to antibiotic-containing medium. Transformed shoots are selected after rooting in medium containing the appropriate antibiotic, transferred to soil and regenerated.

Another aspect of the present invention provides transgenic plants or progeny of these plants containing the isolated DNA of the invention. Both

- 20 monocotyledenous and dicotyledenous plants are contemplated. Plant cells are transformed with the isolated DNA encoding Δ6-desaturase by any of the plant transformation methods described above. The transformed plant cell, usually in a callus culture or leaf disk, is regenerated into a complete transgenic plant by methods well-known to one of ordinary skill in the art (e.g.
- Horsch et al. (1985) Science 227, 1129). In a preferred embodiment, the transgenic plant is sunflower, oil seed rape, maize, tobacco, peanut or soybean. Since progeny of transformed plants inherit the DNA encoding A6-
- desaturase, seeds or cuttings from transformed plants are used to maintain the transgenic plant line.

The present invention further provides a method for providing transgenic plants with an increased content of GLA. This method includes introducing DNA encoding Δ6-desaturase into plant cells which lack or have low levels of GLA but contain LA, and regenerating plants with increased GLA content from the transgenic cells. In particular, commercially grown crop plants are contemplated as the transgenic organism, including, but not limited to, sunflower, soybean, oil seed rape, maize, peanut and tobacco.

The present invention further provides a method for providing transgenic organisms which contain GLA. This method comprises introducing DNA encoding A6desaturase into an organism which lacks or has low levels of GLA, but contains LA. In another embodiment, 15 the method comprises introducing one or more expression vectors which comprise DNA encoding A12-desaturase and Δ6-desaturase into organisms which are deficient in both GLA and LA. Accordingly, organisms deficient in both LA and GLA are induced to produce LA by the expression of A12-desaturase, and GLA is then generated due to the expression of &6-desaturase. Expression vectors comprising DNA encoding \$12-desaturase, or \$12desaturase and \$6-desaturase, can be constructed by methods of recombinant technology known to one of ordinary skill in the art (Sambrook et al., 1989) and the published sequence of \$\textit{al2}\$-desaturase (Wada et al [1990] Nature (London) 347, 200-203. In addition, it has been discovered in accordance with the present 30 invention that nucleotides 2002-3081 of SEQ. ID NO:1 encode cyanobacterial Al2-desaturase. Accordingly, this sequence can be used to construct the subject expression

vectors. In particular, commercially grown crop plants
are contemplated as the transgenic organism, including,
but not limited to, sunflower, soybean, oil seed rape,
maize, peanut and tobacco.

The present invention is further directed to a 5 method of inducing chilling tolerance in plants. Chilling sensitivity may be due to phase transition of lipids in cell membranes. Phase transition temperature depends upon the degree of unsaturation of fatty acids in membrane lipids, and thus increasing the degree of 10 unsaturation, for example by introducing A6-desaturase to convert LA to GLA, can induce or improve chilling resistance. Accordingly, the present method comprises introducing DNA encoding A6-desaturase into a plant cell, and regenerating a plant with improved chilling 15 resistance from said transformed plant cell. In a preferred embodiment, the plant is a sunflower, soybean, oil seed rape, maize, peanut or tobacco plant.

The following examples further illustrate the present invention.

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EXAMPLE 1

Strains and Culture Conditions

Synechocystis (PCC 6803, ATCC 27184), Anabaena (PCC 7120, ATCC 27893) and Synechococcus (PCC 7942, ATCC 33912) were grown photoautotrophically at 30°C in BG11N+medium (Rippka et al. [1979] J. Gen. Microbiol. 111, 1-61) under illumination of incandescent lamps (60µE.m⁻².S⁻¹). Cosmids and plasmids were selected and propagated in Escherichia coli strain DH5a on LB medium supplemented with antibiotics at standard concentrations as described by Maniatis et al. (1982) Molecular

Cloning: A Laboratory Manual, Cold Spring Harbor

Laboratory, Cold Spring, New York.

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EXAMPLE 2 ı

Construction of Synechocystis Cosmid Genomic Library Total genomic DNA from Synechocystis (PCC 6803) was partially digested with Sau3A and fractionated on a sucrose gradient (Ausubel et al. [1987] Current Protocols in Molecular Biology, Greene Publishing Associates and Wiley Interscience, New York). Fractions containing 30 to 40 kb DNA fragments were selected and ligated into the dephosphorylated BamHI site of the cosmid vector, pDUCA7 (Buikema et al. [1991] J. Bacteriol. 173, 1879-1885). The ligated DNA was packaged in vitro as described by Ausubel et al. (1987), and packaged phage were propagated in E. coli DH5a containing the AvaI and Eco4711 methylase helper plasmid, pRL528 as described by Buikema et al. (1991). 15 A total of 1152 colonies were isolated randomly and maintained individually in twelve 96-well microtiter plates.

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EXAMPLE 3

Gain-of-Function Expression of GLA in Anabaena Anabaena (PCC 7120), a filamentous cyanobacterium, is deficient in GLA but contains significant amounts of linoleic acid, the precursor for GLA (Figure 2; Table 2). The Synechocystis cosmid library described in Example 2 was conjugated into Anabaena (PCC 7120) to identify transconjugants that produce GLA. Anabaena cells were grown to mid-log phase in BG11N+ liquid medium and resuspended in the same medium to a final concentration of approximately 2x10° cells per ml. A mid-log phase culture of E. coli RP4 (Burkardt et al. [1979] J. Gen. Microbiol. 114, 341-348) grown in LB containing ampicillin was washed and resuspended in fresh LB medium. Anabaena and RP4 were then mixed and spread evenly on BG11N+ plates containing The cosmid genomic library was replica plated onto LB plates containing 50 µg/ml kanamycin and 17.5 μg/ml chloramphenicol and was subsequently patched onto BG11N+ plates containing Anabaena and RP4. After 24 hours of incubation at 30°C, 30 µg/ml of neomycin was underlaid; and incubation at 30°C was continued until transconjugants appeared.

Individual transconjugants were isolated after
conjugation and grown in 2 ml BG11N+ liquid medium with
15 µg/ml neomycin. Fatty acid methyl esters were
prepared from wild type cultures and cultures containing
pools of ten transconjugants as follows. Wild type and
transgenic cyanobacterial cultures were harvested by
centrifugation and washed twice with distilled water.
Fatty acid methyl esters were extracted from these
cultures as described by Dahmer et al. (1989) J. Amer.

- Oil. Chem. Soc. 66, 543-548 and were analyzed by Gas
 Liquid Chromatography (GLC) using a Tracor-560 equipped
 with a hydrogen flame ionization detector and capillary
 column (30 m x 0.25 mm bonded FSOT Superox II, Alltech
- 5 Associates Inc., IL). Retention times and cochromatography of standards (obtained from Sigma
 Chemical Co.) were used for identification of fatty
 acids. The average fatty acid composition was
 determined as the ratio of peak area of each C18 fatty
 10 acid normalized to an internal standard.

Representative GLC profiles are shown in Fig. 2. C18 fatty acid methyl esters are shown. Peaks were identified by comparing the elution times with known standards of fatty acid methyl esters and were confirmed

- by gas chromatography-mass spectrometry. Panel A depicts GLC analysis of fatty acids of wild type Anabaena. The arrow indicates the migration time of GLA. Panel B is a GLC profile of fatty acids of transconjugants of Anabaena with pAM542+1.8F. Two GLA
- producing pools (of 25 pools representing 250 transconjugants) were identified that produced GLA. Individual transconjugants of each GLA positive pool were analyzed for GLA production; two independent transconjugants, AS13 and AS75, one from each pool, were
- identified which expressed significant levels of GLA and which contained cosmids, cSy13 and cSy75, respectively (Figure 3). The cosmids overlap in a region approximately 7.5 kb in length. A 3.5 kb NheI fragment of cSy75 was recloned in the vector pDUCA7 and
- 30 transferred to <u>Anabaena</u> resulting in gain-of-function expression of GLA (Table 2).

- Two Nhel/Hind III subfragments (1.8 and 1.7 kb) 1 of the 3.5 kb Nhe I fragment of cSy75-3.5 were subcloned into "pBLUESCRIPT" (Stratagene) (Figure 3) for sequencing. Standard molecular biology techniques were performed as described by Maniatis et al. (1982) and Ausubel et al. (1987). Dideoxy sequencing (Sanger et al. [1977] Proc. Natl. Acad. Sci. USA 74, 5463-5467) of pBS1.8 was performed with "SEQUENASE" (United States Biochemical) on both strands by using specific oligonucleotide primers synthesized by the Advanced DNA 10 Technologies Laboratory (Biology Department, Texas A & M University). DNA sequence analysis was done with the GCG (Madison, WI) software as described by Devereux et al. (1984) Nucleic Acids Res. 12, 387-395.
- Both NheI/HindIII subfragments were transferred into a conjugal expression vector, AM542, in both forward and reverse orientations with respect to a cyanobacterial carboxylase promoter and were introduced into Anabaena by conjugation. Transconjugants

 20 containing the 1.8 kb fragment in the forward orientation (AM542-1.8F) produced significant quantities of GLA and octadecatetraenoic acid (Figure 2; Table 2). Transconjugants containing other constructs, either reverse oriented 1.8 kb fragment or forward and reverse oriented 1.7 kb fragment, did not produce detectable levels of GLA (Table 2).

Figure 2 compares the C18 fatty acid profile of an extract from wild type Anabaena (Figure 2A) with that of transgenic Anabaena containing the 1.8 kb fragment of cSy75-3.5 in the forward orientation (Figure 2B). GLC analysis of fatty acid methyl esters from AM542-1.8F revealed a peak with a retention time identical to that

1	of authentic GLA standard. Analysis of this peak by gas
	chromatography-mass spectrometry (GC-MS) confirmed that
	it had the same mass fragmentation pattern as a GLA
	reference sample. Transgenic Anabaena with altered
5	levels of polyunsaturated fatty acids were similar to
	wild type in growth rate and morphology.

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Table 2
Composition of C18 Fatty Acids in
Wild Type and Trasgenic Cyanobacteria

5	Strain		F	atty a	cid (%)								
		18:0	18:1	18:2	18:3 (a)	18:3 (γ)	. 18:4							
	Wild type													
10	Synechocystis (sp.PCC6803)	13.6	4.5	54.5	_	27.3	-							
10	Anabaena (sp.PCC7120)	2.9	24.8	37.1	35.2 -		-							
	Synechococcus (Sp.PCC7942)	20.6	79.4	-	-	-								
	Anabaena Transconjugants													
15	cSy75	3.8	24.4	22.3	9.1	27.9	12.5							
	cSy75-3.5	4.3	27.6	18.1	3.2	40.4	6.4							
	pΛM542-1.8F	4.2	13.9	12.1	19.1	25.4	25.4							
	pΛM542-1.8R	7.7	23.1	38.4	30.8	-	-							
20	pΛM542-1.7F	2.8	27.8	36.1	33.3	-	-							
	pΛM542-1.7R	2.8	25.4	42.3	29.6	-	-							
	Synechococcus Transformants													
	ρΛM854	27.8	72.2	-	-	1								
	pΛM854-Δ ¹²	4.0	43.2	46.0	_	-	J - 1							
25	рЛМ854-∆ €	18.2	81.8	-	_	-	-							
	pΛM854-Δ ⁶ & Δ ¹²	42.7	25.3	19.5	-	16.5	-							

^{18:0,} stearic acid; 18.1, oleic acid; 18:2, linoleic acid; 18:3(α), α-linolenic acid; 18:3(γ), γ-linolenic acid; 18:4, octadecatetraenoic acid

1 EXAMPLE 4

Transformation of Synechococcus with A6 and A12 Desaturase Genes

A third cosmid, cSy7, which contains a \$12desaturase gene, was isolated by screening the 5 Synechocystis genomic library with a oligonucleotide synthesized from the published Synechocystis A12desaturase gene sequence (Wada et al. [1990] Nature (London) 347, 200-203). A 1.7 kb AvaI fragment from this cosmid containing the A12-desaturase gene was 10 identified and used as a probe to demonstrate that cSy13 not only contains a A6-desaturase gene but also a A12desaturase gene (Figure 3). Genomic Southern blot analysis further showed that both the \$6-and \$12desaturase genes are unique in the Synechocystis genome 15 so that both functional genes involved in C18 fatty acid desaturation are linked closely in the Synechocystis genome.

The unicellular cyanobacterium Synechococcus (PCC 7942) is deficient in both linoleic acid and GLA(3).

The \$\textit{\alpha}\$12 and \$\textit{\alpha}\$6-desaturase genes were cloned individually and together into pAM854 (Bustos et al. [1991] \$\frac{\textit{J.}}{\textit{Bacteriol.}}\$ \frac{174}{\textit{7525-7533}}\$, a shuttle vector that contains sequences necessary for the integration of foreign DNA into the genome of \$\frac{\text{Synechococcus}}{\text{al.}}\$ (Golden et al. [1987] \$\frac{\text{Methods in Enzymol.}}{\text{153}}\$, 215-231).

Synechococcus was transformed with these gene constructs and colonies were selected. Fatty acid methyl esters were extracted from transgenic \$\frac{\text{Synechococcus}}{\text{and}}\$ analyzed by GLC.

Table 2 shows that the principal fatty acids of wild type Synechococcus are stearic acid (18:0) and

- l oleic acid (18:1). <u>Synechococcus</u> transformed with pAM854-Δ12 expressed linoleic acid (18:2) in addition to the principal fatty acids. Transformants with pAM854-Δ6 and Δ12 produced both linoleate and GLA (Table 1).
- 5 These results indicated that Synechococcus containing both \$12-\$ and \$6-\$desaturase genes has gained the capability of introducing a second double bond at the \$12 position and a third double bond at the \$6 position of C18 fatty acids. However, no changes in fatty acid composition was observed in the transformant containing
- composition was observed in the transformant containing pAM854-\$\(\alpha \), indicating that in the absence of substrate synthesized by the \$\(\alpha \)12 desaturase, the \$\(\alpha \)6-desaturase is inactive. This experiment further confirms that the 1.8 kb \(\alpha \)heI/HindIII fragment (Figure 3) contains both coding
- and promoter regions of the <u>Synechocystis</u> \$46-desaturase gene. Transgenic <u>Synechococcus</u> with altered levels of polyunsaturated fatty acids were similar to wild type in growth rate and morphology.

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1 EXAMPLE 5

Nucleotide Sequence of A6-Desaturase

The nucleotide sequence of the 1.8 kb fragment of cSy75-3.5 including the functional \$\triangle 6\$-desaturase gene

5 was determined. An open reading frame encoding a polypeptide of 359 amino acids was identified (Figure 4). A Kyte-Doolittle hydropathy analysis (Kyte et al. [1982] J. Mol. Biol. 157, 105-132) identified two regions of hydrophobic amino acids that could represent transmembrane domains (Figure 1A); furthermore, the hydropathic profile of the \$\triangle 6\$-desaturase is similar to that of the \$\triangle 12\$-desaturase gene (Figure 1B; Wada et al.) and \$\triangle 9\$-desaturases (Thiede et al. [1986] J. Biol. Chem. 261, 13230-13235). However, the sequence similarity between the Synechocystis \$\triangle 6\$- and \$\triangle 12\$-desaturases is less than 40% at the nucleotide level and approximately 18% at the amino acid level.

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1 EXAMPLE 6

Transfer of Cyanobacterial & -Desaturase into Tobacco The cyanobacterial ^6-desaturase gene was mobilized into a plant expression vector and transferred to tobacco using Agrobacterium mediated gene transfer To ensure that the transferred desaturase techniques. is appropriately expressed in leaves and developing seeds and that the desaturase gene product is targeted to the endoplasmic reticulum or the chloroplast, various 10 expression cassettes with Synechocystis &-desaturase open reading frame (ORF) were constructed. Components of these cassettes include: (i) a 35S promoter or seed specific promoter derived from the sunflower helianthinin gene to drive \$\delta^6\$-desaturase gene expression 15 in all plant tissues or only in developing seeds respectively, (ii) a putative signal peptide either from carrot extensin gene or sunflower helianthinin gene to target newly synthesized Δ^6 -desaturase into the ER, (iii) an ER lumen retention signal sequence (KDEL) at 20 the COOH-terminal of the Δ^6 -desaturase ORF, and (iv) an optimized transit peptide to target \$\delta^6\$ desaturase into the chloroplast. The 35S promoter is a derivative of pRTL2 described by Restrepo et al. (1990). optimized transit peptide sequence is described by Van 25 de Broeck et al. (1985). The carrot extensin signal peptide is described by Chen et al (1985) EMBO J. 9, 2145.

Transgenic tobacco plants were produced containing a chimeric cyanobacterial desaturase gene, 30 comprised of the <u>Synechocystis</u> Δ^6 desaturase gene fused to an endoplasmic reticulum retention sequence (KDEL) and extensin signal peptide driven by the CaMV 35s promoter. PCR amplifications of transgenic tobacco genomic DNA indicate that the Δ^6 desaturase gene was incorporated into the tobacco genome. Fatty acid methyl esters of leaves of these transgenic tobacco plants were

1	extracted and analyzed by Gas Liquid Chromatography
	(GLC). These transgenic tobacco accumulated significant
	amounts of GLA (Figure 4). Figure 4 shows fatty acid
	methyl esters as determined by GLC. Peaks were
5	identified by comparing the elution times with known
	standards of fatty acid methyl ester. Accordingly,
,	cyanobacterial genes involved in fatty acid metabolism
1	can be used to generate transgenic plants with altered

fatty acid compositions.

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1	SEQUENCE	LISTING
	(1) GENERAL INFORMATION:	
5	(i) APPLICANT: Thomas, Terry L. Reddy, Avutu S. Nuccio, Michael Freyssinet, Georges	L.
	(ii) TITLE OF INVENTION: PRODUCTION ACID BY A I	OF GAMMA LINOLENIC DELTA 6-DESATURASE
10	(iii) NUMBER OF SEQUENCES: 3	
15	<pre>(iv) CORRESPONDENCE ADDRESS: (A) ADDRESSEE: Scully, Scott, (B) STREET: 400 Garden City PI (C) CITY: Garden City (D) STATE: New York (E) COUNTRY: United States (F) ZIP: 11530</pre>	Murphy & Presser Laza
	(v) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatibl (C) OPERATING SYSTEM: PC-DOS/N (D) SOFTWARE: Patentin Release	IS-DOS
20 .	(vi) CURRENT APPLICATION DATA: (A) APPLICATION NUMBER: To be (B) FILING DATE: 08-JAN-1992 (C) CLASSIFICATION:	e assigned
25	<pre>(viii) ATTORNEY/AGENT INFORMATION: (A) NAME: McNulty, William E. (B) REGISTRATION NUMBER: 22,60 (C) REFERENCE/DOCKET NUMBER: 8</pre>	96 3383Z
	(ix) TELECOMMUNICATION INFORMATION: (A) TELEPHONE: (516) 742-4343 (B) TELEFAX: (516) 742-4366 (C) TELEX: 230 901 SANS UR	
30	•	

1	(2) INFORMATION FOR SEQ ID NO:1:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 3588 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
1	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 20023081	
1	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
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	CACCTTGCCA GACCACGTTA GTTTGAGTGT TTCCGCCCTG GCGGCCCCGA TTTTTTCCTT	18
	TGCGGCTTTG GGCAATCAGG CGATCGGGCA ATTGCGTTTG TTTGACCAGA CTTGGCCCAT	240
	TCAGGAAATT GTCATTCACC AAGACCATCC CTGGCTCAAT TTACCCCTGG CGGATTTATG	30
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	AGGCGCAGTG GTGAATAATT TAACGTTGCA ATCTGGGGAC CATTTAATAG TGGGACAAAA	42
	ACCCCAACCC AAGACCAAAC GGCGATCGCC TTGGCGCAAA TTTTCCAAAC TGATTACCAA	48
	CCTGCGGGAG TATCAGCGGT ATGTCCAACA GGTGATATGG GTGGTGTTGT TITTATTGTT	54
	GATGATITT CTGGCCACCT TCATCTACGT TTCCATTGAT CAACATATTG CCCCAGTGGA	60
20	CGCGTTGTAT TTTTCCGTGG GCATGATTAC CGGGGCCGGT GGCAAGGAAG AGGTGGCCGA	66
	AAAGTCCCCC GATATCATCA AAGTATTCAC AGTGGTGATG ATGATCGCCG GGGCGGGGT	72
	GATTGGTATT TGTTATGCCC TACTGAATGA TTTCATCCTT GGCAGTCGCT TTAGTCAGTT	78
	TTTGGATGCG GCCAAGTTAC CCGATCGCCA TCACATCATC ATTTGTGGGC TGGGGGGAGT	84
25	GAGCATGGCC ATTATTGAAG AGTTAATTCA CCAGGGCCAT GAAATTGTGG TAATCGAAAA	90

GGATACAGAT AATCGTTTCT TGCATACGGC CCGCTCCCTG GGGGTGCCCG TAATTGTGGA

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•	CCCTAGCCTG	CCAGTGGTGT	TGCGTTGCCA	GGATGCCCAG	TTTAGCCTGT	CCCTGCAGGA	1140
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. 5					GATGATTTGC		1260
•					CAATTGGTTA		1320
					GGCAAAACCA	•	1380
					TATTTAACCA		1440
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. 10					GGTGATCAAG		1560
					GATAACTTCC		1620
				•	GTTTTACTGC		1680
-	TGCAAAAAAG						1740
15				•	AGTATTTTCT		1800
					CCAATATTGC		1860
					ACTCCCAGTT		1920
					GCGGTATAAT		1980
					AGA ATT AAA		. 2031
20			Met Leu 7 1	Chr Ala Glu	Arg Ile Lys	Phe Thr	. 2031
	CAG AAA CGG	GGG TTT CG	T CGG GTA C	TA AAC CAA	CGG GTG GAT		2079
	Gln Lys Arg	Gly Phe Ar 15	g Arg Val I	eu Asn Gln 20	Arg Val Asp	Ala Tyr	2019
	TTT GCC GAG	CAT GGC CT	G ACC CAA A	GG GAT AAT	CCC TCC ATG	mam eme	2127
	Phe Ala Glu	His Gly Le	u Thr Gln A	rg Asp Asn 35	Pro Ser Met	Tyr Leu	2121
25							

1			ATT											2175
_			CCA Pro											2223
5			GCC Ala											2271
			GCC Ala											2319
10			GAT Asp 110											2367
			TTG Leu											2415
15			GGA Gly											2463
			TAT Tyr											2511
			TTT Phe											2559
20			TAT Tyr 190					_						2607
			TTG Leu											2655
25		Pro	CTG Leu	Ala	Leu	Gly	Phe	Ser	Ile	Pro	Glu	Val		2703

~																	
1	GCT Ala 235	Set	GTA Val	ACC	TAT Tyr	ATG Met 240	ACC Thr	TAT Tyr	GGC Gly	Ile	GTG Val 245	GTT Val	TGC Cys	ACC Thr	ATC Ile	TIT Phe 250	2751
5	ATG Met	CTG Leu	GCC Ala	CAT His	GTG Val 255	TTG Leu	GAA Glu	TCA Ser	ACT Thr	GAA Glu 260	TTT Phe	CTC Leu	ACC Thr	CCC Pro	GAT Asp 265	GGT Gly	2799
)	GAA Glu	TCC Ser	GGT Gly	GCC Ala 270	ATT Ile	GAT Asp	GAC Asp	GAG Glu	TGG Trp 275	GCT Ala	ATT	TGC Cys	CAA Gln	ATT Ile 280	CGT Arg	ACC Thr	2847
	ACG Thr	GCC Ala	AAT Asn 285	TTT Phe	GCC Ala	ACC Thr	AAT Asn	AAT Asn 290	CCC Pro	TTT Phe	TGG Trp	AAC Asn	TGG Trp 295	TTT Phe	TGT Cys	GGC Gly	2895
10	GGT Gly	TTA Leu 300	AAT Asn	CAC His	CAA Gln	GTT Val	ACC Thr 305	CAC His	CAT His	CTT Leu	TTC Phe	CCC Pro 310	AAT Asn	ATT Ile	TGT Cys	CAT His	2943
	ATT Ile 315	CAC His	TAT Tyr	CCC Pro	CAA Gln	TTG Leu 320	GAA Glu	AAT Asn	ATT Ile	ATT Ile	AAG Lys 325	GAT Asp	GTT Val	TGC Cys	CAA Gln	GAG Glu 330	2991
15	TTT Phe	GGT Gly	GTG Val	GAA Glu	TAT Tyr 335	AAA Lys	GTT Val	TAT Tyr	CCC Pro	ACC Thr 340	Phę	AAA Lys	GCG Ala	GCG Ala	ATC Ile 345	GCC Ala	3039
	TCT Ser	AAC Asn	TAT Tyr	CGC Arg 350	TGG Trp	CTA Leu	GAG Glu	GCC Ala	ATG Met 355	GGC Gly	AAA Lys	GCA Ala	TCG Ser	TGAC	ATTG	scc .	3088
	TTG	GATT	GA A	AGCAA	TAAL	G CA	L AAA	CCCI	CGI	'AAAT	CTA	TGAT	CGAA	GC C	TTTC	TGTTG	3148
00	ccc	SCCG#	CC. Y	AATC	ccce	A TO	CTGA	CCAA	AGG	TTGA	TĢT	TGGC	ATTG	CT C	CAAA	CCCAC	3208
50																TGATT	3268
																CTCAA	3328
																CCATG	3388
	TGGI	CTAA	cc c	AGCC	CTGG	C CA	AGGC	TTGG	AÇC	AAGG	CCA	TGCA	TTAA	CT C	CACG	AGGCT	3448
25	AGGC	CAGA	AA A	ATTA	TAT.	'G GC	TCCT	'GATI	TCT	TCCG	GCT	ATCG	CACC	TA C	CGAT	TITTG	3508
-	AGCA	TTT	TG C	CAAG	GAAI	т ст	ATCC	CCAC	TAT	CTCC	ATC	CCAC	TCCC	CC G	ССТС	TACAA	3568

-34-

1	AAT	TTTA'	TCC	ATCA	GCTA	GC							•				3588
•	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:2	:								
5			(i)	(A (B	ENCE) LEI) TY:) TO:	NGTH PE:	: 35 amin	9 am.	ino a id		5						
		(ii)	MOLE	CULE	TYP	E: p	rote	in								
! !		(:	xi)	SEQU	ENCE	DES	CRIP'	rion	: SE	QI Q	NO:	2:					
/	Met 1	Leu	Thr	Ala	Glu 5	Arg	Ile	Lys	Phe	Thr 10	Gln	Lys	Arg	Gly	Phe 15	Arg	
10	Arg	Val	Leu	Asn 20	Gln	Arg	Val	Asp	Ala 25	Tyr	Phe	Ala	Glu	His 30	Gly	Leu	•
	Thr	Gln	Arg 35	Asp	Asn	Pro	Ser	Met 40	Tyr	Leu	Lys	Thr	Leu 45	Ile	Ile	Val	
	Leu	Trp 50	Leu	Phe	Ser	Ala	Trp 55	Ala	Phe	Val	Leu	Phe 60	Ala	Pro	Val	Ile	
15	Phe 65	Pro	Val	Arg	Leu	Leu 70	Gly	Cys	Met	Val	Leu 75	Ala	Ile	Ala	Leu	Ala 80	
	Ala	Phe	Ser	Phe	Asn 85	Val	Gly	His	Asp	Ala 90	Asn	His	Asn	Ala	Tyr 95	Ser	
	Ser	Asn	Pro	His 100	Ile	Asn	Arg	Val	Leu 105	Gly	Met	Thr	Tyr	Asp 110	Phe	Val	
20	Gly	Leu	Ser 115	Ser	Phe	Leu	Trp.	Arg 120	Tyr	Arg	His	Asn	Tyr 125	Leu	His	His	
	Thr	Tyr 130	Thr	Asn	Ile	Leu	Gly 135	His	Asp	Val	Glu	Ile 140	His	Gly	Asp	Gly	
	Ala 145	Val	Arg	Met	Ser	Pro 150	Glu	Gln	Glu	His	Val 155	Gly	Ile	Tyr	Arg	Phe 160	
25	Gln	Gln	Phe	Tyr	Ile 165	Trp	Gly	Leu	Tyr	Leu 170	Phe	Ile	Pro	Phe	Tyr 175	Trp	

30

_																
1				100				·Val	182		•			190		
'	His	Lys	Ile 195	Pro	Pro	Phe	Gln	Pro 200	Leu	Glu	Leu	Ala	Ser 205	Leu	Leu	Gly
5	Ile	Lys 210	Leu	Leu	Trp	Leu	Gly 215	Tyr	Val	Phe	Gly	Leu 220	Pro	Leu	Ala	Leu
	Gly 225	Phe	Ser	Ile	Pro	Glu 230	Val	Leu	Ile	Gly	Ala 235	Ser	Val	Thr	Tyr	Met 240
								Thr		250		•			255	
	Glu	Ser	Thr	Glu 260	Phe	Leu	Thr	Pro	Asp 265	Gly	Glu	Ser	Gly	Ala 270	Île	Asp
								Ile 280					285	•		
	Asn	Asn 290	Pro	Phe	Trp	Asn	Trp 295	Phe	Cys	Gly	Gly	Leu 300	Asn	His	Gln	Val
15	Thr 305	His	His	Leu	Phe	Pro 310	Asn	Ile	Cys	His	Ile 315	His	Tyr	Pro	Gln	Leu 320
ر ــــ	Glu	Asn	Ile	Ile	Lys 325	Asp	Val	Cys	Gln	Glu 330	Phe	Gly	Val	Glu	Tyr 335	Lys
	Val	Tyr	Pro	Thr 340	Phe	Lys	Ala	Ala	11e 345	Ala	Ser	Asn	Tyr	Arg 350	Trp	Leu
	Glu	Ala	Met 355	Gly	Lys	Ala	Ser									
20	(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	0:3:		•						
		(i)	(A (B (C) LE) TY) ST	NGTH PE: RAND	ARAC : 18 nucl EDNE GY:	84 b eic SS:	both	S: pair	s			-			٠

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(ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

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1	AGCTTCACTT	CGGTTTTATA	TTGTGACCAT	GGTTCCCAGG	CATCTGCTCT	AGGGAGTTTT	. 60
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10	CCAGTTATTT	TTCCGGTGCG	CCTACTGGGT	TGTATGGTTT	TGGCGATCGC	CTTGGCGGCC	540
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	GGCTAGGCCA	GAAAAATTAT	ATTGGCTCCT	GATTTCTTCC	GGCTATCGCA	CCTACCGATT	1800
	TTTGAGCATT	TTTGCCAAGG	AATTCTATCC	CCACTATCTC	CATCCCACTC	сссесстет	1860
	ACAAAATTTT	ATCCATCAGC	TAGC		•		1884

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1 WHAT IS CLAIMED:

- 1. An isolated nucleic acid encoding bacterial \$46-desaturase.
- 2. The nucleic acid of Claim 1 comprising the 5 nucleotides of SEQ. ID NO:3.
 - 3. An isolated nucleic acid that codes for the amino acid sequence encoded by the nucleic acid of Claim 1.
 - 4. The isolated nucleic acid of any one of Claims 1-3 wherein said nucleic acid is contained in a vector.
- 10 5. The isolated nucleic acid of Claim 4 operably linked to a promoter and/or a termination signal capable of effecting expression of the gene product of said isolated nucleic acid.
- 6. The isolated nucleic acid of Claim 5 wherein said promoter is a \$\(^6\)-desaturase promoter, an Anabaena carboxylase promoter, a helianthinin promoter, a glycin promoter, a napin promoter, or a helianthinin tissue-specific promoter.
- The isolated nucleic acid of Claim 5 wherein said termination signal is a <u>Synechocystis</u> termination signal, a nopaline synthase termination signal, or a seed termination signal.
 - 8. The isolated nucleic acid of any one of Claims 1-7 wherein said isolated nucleic acid is contained within a transgenic organism.
- 9. The isolated nucleic acid of Claim 8 wherein said transgenic organism is a bacterium, a fungus, a plant cell or an animal.
 - 10. A plant or progeny of said plant which has been regenerated from the transgenic plant cell of Claim 9.
- 30 11. The plant of Claim 10 wherein said plant is a sunflower, soybean, maize, tobacco, peanut or oil seed rape plant.

- 1 12. A method of producing a plant with increased gamma linolenic acid (GLA) content which comprises:
 - (a) transforming a plant cell with the isolated nucleic acid of any one of Claims 1-7; and
- 5 (b) regenerating a plant with increased GLA content from said plant cell.
 - 13. The method of Claim 12 wherein said plant is a sunflower, soybean, maize, tobacco, peanut or oil seed rape plant.
- 14. A method of inducing production of gamma linolenic acid (GLA) in an organism deficient or lacking in GLA with comprises transforming said organism with the isolated nucleic acid of any one of Claims 1-7.
- 15. A method of inducing production of gamma
 15 linolenic acid (GLA) in an organism deficient or lacking in
 GLA and linoleic acid (LA) which comprises transforming said
 organism with an isolated nucleic acid encoding bacterial \$\times 6\$desaturase and an isolated nucleic acid encoding \$\times 12\$desaturase.
- 20 16. A method of inducing production of gamma linolenic acid (GLA) in an organism deficient or lacking in GLA and linoleic acid (LA) which comprises transforming said organism with at least one expression vector comprising an isolated nucleic acid encoding bacterial \$\delta\$6-desaturase and an isolated nucleic acid encoding \$\delta\$12-desaturase.
 - 17. The method of any one of Claims 15 or 16 wherein said isolated nucleic acid encoding $\Delta 6$ -desaturase comprises nucleotides 317 to 1507 of SEQ. ID NO:1.
- 18. A method of inducing production of
 30 octadecatetraeonic acid in an organism deficient or lacking
 in gamma linolenic acid with comprises transforming said
 organism with isolated nucleic acid of any one of Claims 1-7.

WO 93/06712 PCT/US92/08746

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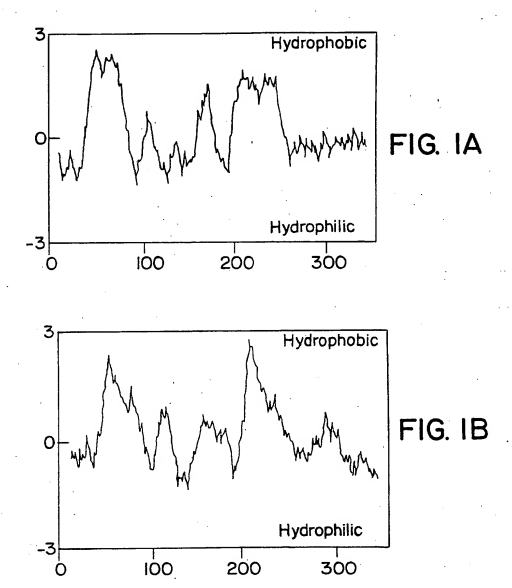
1	19.	The method	of	Claim	18	wherein	said	organism	is	а
	bacterium.	a fungus, a	pla	ant or	an	animal.				

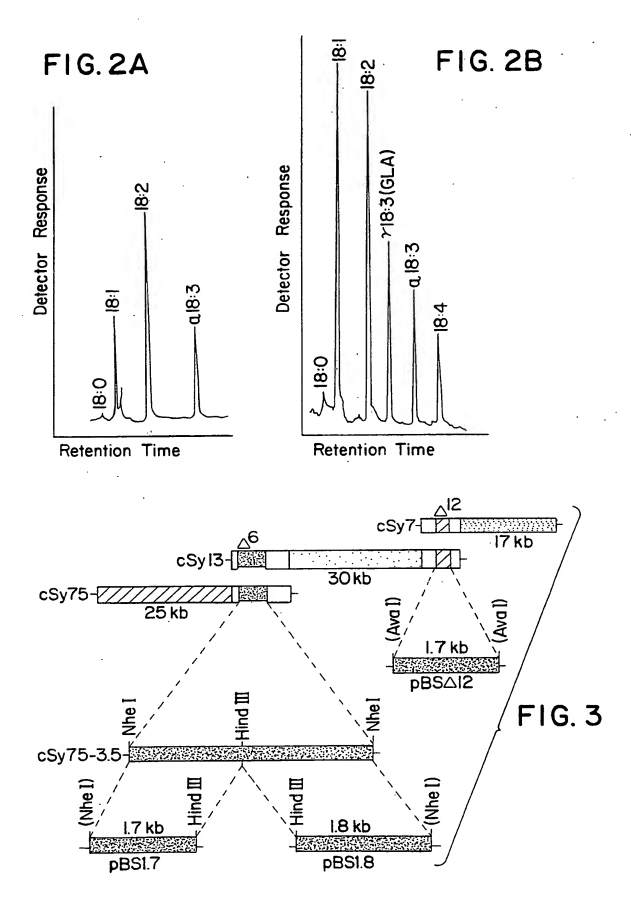
- 20. A method of use of the isolated nucleic acid of any one of Claims 1-7 to produce a plant with improved5 chilling resistance which comprises:
 - a) transforming a plant cell with the isolated nucleic acid of any one of Claims 1-7; and
 - b) regenerating said plant with improved chilling resistance from said transformed plant cell.
- 21. The method of Claim 20 wherein said plant is a sunflower, soybean, maize, tobacco, peanut or oil seed rape plant.
 - 22. Isolated bacterial A6-desaturase.
- 23. The isolated bacterial $_{\Delta}6$ -desaturase of Claim 22 15 which has an amino acid sequence of SEQ ID NO:2.

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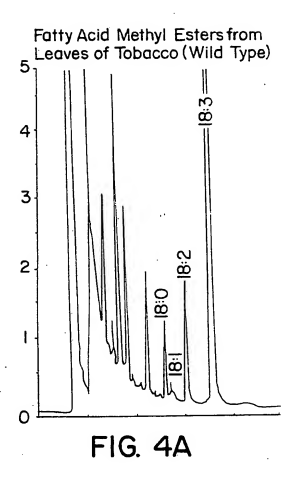
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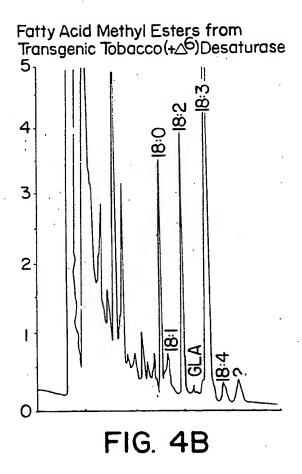
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SUBSTITUTE SHEET





INTERNATIONAL SEARCH REPORT

International Application No. PCT/US92/08746

A. CLASSIFICATION OF SUBJECT MATTER IPC(5) :Please See Extra Sheet. US CL. :800/205; 455/172.3, 189, 69.1, 320.1, 134, 170, 171; 536/27 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) U.S. : 800/205; 435/172.3, 189, 69.1, 320.1, 134, 170, 171; 536/27; 935/9, 30, 6, 24, 29, 38 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) STN/BIOSIS, CA; APS search terms: inclonic, desaturase, delta-6, gene, DNA, cDNA, puriff, cyanobacteri? C. DOCUMENTS CONSIDERED TO BE RELEVANT Category* Clation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Y. Nature, Volume 347, issued 13 September 1990, H. Wade et al., "Enhancement of Chilling Tolerance of a Cyanobacterium by Genetic Manipulation of Fatty Acid Desaturation", pages 200-203, especially pages 201-203. Y. Biochemical Journal, Volume 240, issued 1986, S. Stymne et al., "Biosynthesis of y-Uninotenic Acid in Cotyledons and Microsomal Preparations of the Developing Seculs of Common Borage (Borago officinalis)", pages 385-392, especially pages 385 and 392. P.P. A., 0.255, 378 (Kridl et al.) 3 February 1988, see entire document, especially columns 3-5 and 7-11. See patent far tilly annex. * Special estageries of clied documents: * * * Special estageries of clied documents: * * * Special estageries of clied documents: * * * * Special estageries of clied docu					
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cited to establish the publication date of another citation or other special reason (as specified) Y document of particular relevance; the claimed invention cannot be	cited to establish the publication date of another citation or other	"Y" document of particular relevance; the claimed invention cannot be			
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Date of the actual completion of the international search Date of mailing of the international search 1 2 IAN 1003	Date of the actual completion of the international search 03 DECEMBER 1992	Date of mailing of the international search report 13 JAN 1993			
03 DECEMBER 1992	Name and mailing address of the ISA/ Commissioner of Patents and Trademarks	Authorized officer			
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INTERNATIONAL SEARCH REPORT

International Application No. PCT/US92/08746

A. CLASSIFICATION OF SUBJECT MATTER: IPC (5):		
A01H 1/00, 5/00; C12N 15/00, 9/02; C12P 7/64, 1/02, 1/0	4, 21/06; C07H 15/12, 17/	00
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